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(54) Resin-reinforced elastomer, process for producing same and pneumatic tire using same

(57) There are disclosed a resin-reinforced elestomer which comprises as principal components, a rubbra and a polyoletin having an average particle diameter of at most 1 µm, and in which the rubber and the polyoletin are linked together, a processe for producing the resin-reinforced elastomer by metit kneading a polyolefin, a No.1 rubbor and shinding agent to prepare a thermoplastic composition, adding a No.2 rubber to the resultant thermoplastic composition and malt kneading the mixture thus formed to disperse the polyolefin in the rubber components; and a pneumatic tire using the resin-teniforced elastomer in its tread. The above resin-teniforced elastomer in its tread. The above resin-teniforced elastomer has a uniform modulus and a low density, is minimized in directionality, and is excellent in teneile strength, fetigue resistance and abrasion resistance.

Description

The present invention relates to a resin-reinforced elastomer, a process for producing the same and a pneumatic tire using the same. More particularly, the present invention pertains to a resin-reinforced elastomer which is excellent in modulus, strength, wear resistance, taligue resistance and the like and which has a low density; to a processor of efficiently producing said resin-reinforced elastomer, and to a pneumatic tire in which said resin-reinforced elastomer having the aforesaid characteristics is used in the tread.

A resin-reinforced elastomer, that is, the composition in which a small amount of a polyolellin is dispersed in a vulcanizable rubber such as natural rubber, polyisopraer rubber, polybuddene rubber and ethylene-propylene rubber to improve the crack growth resistance, modulus and strength has heretolore been produced in general by a method in which a rubber is blended with a polyolelfin such as polyethylene, polypropylene and ethylene-propylene copolymer and then the resultant blend is vulcanized.

For example, Japanese Patent Publication No. 78155/1995 (Hei-7) describes a thermoplastic elastomer composition which provides molded articles such as a car bumper that are minimized in dependence of tensile strength on truckness and in anisotropy and have favorable moldebility as well as excellent external-appearance, while maintaining impact resistance.

In addition, Japanese Patent Provisional Publication No. 186606/1995(Hei-7) discloses a pneumatic tire which comprises a rubber composition containing a thermoplastic resin in its tread rubber and which is improved in rolling resistance, white maintaining favorable traction characteristics.

However, the resin-reinforced elastomer obtained by the aforesaid method involves the problems of its having directionally because of the oriented resin, deficiency in strength, wear resistance and falling resistance and the like, when being used in a tire member of an automobile. It is desirable that a tread pubber be excellent in strength, wear resistance, out-through resistance and durability against heat generation, each being well-balanced with one enother, and also be improved in extrusion processability. With regard to the above-mentioned requirements, the conventional resin-reinforced elastomer some mentioned above falls to sufficiently satisfy the required characteristics. Thus the present actual-situation is that the development of a resin-reinforced elastomer fully improved in the above-mentioned problems is exampled in the supervision of the su

A method in which low density polyethylene is blended to form a rubber composition is known as an efective means for enhancing various performances of a tire. Nevertheless, the blending of the low density polyethylene is unfavorable when used in a rubber composition for a tire which is raised in temperature during running, since it has a low mething point and thus remarkably changes the physical properties or the rubber composition while the temperature is elevated. On the other hand, the use of high density polyethylene in place thereof for the purpose of raising the mething point suffers the disadvantage that the Mooney viscosity of the unvulcant-zed rubber composition is raised, thereby causeing extreme difficulty in extrusion processina, as described in USP No. 5.41.88.

Under such circumstances, the object of the present invention is to provide a resin-reinforced elastomer which is excellent in extrusion processability when being used in unvulcanized state, and which is characterized when made into a vulcanizate, by its uniform modulus, being minimized in directionality, being excellent in tensile istrangth, absort resistance and fatigue resistance and having a low density. Another object thereof is to provide a process for producing said resin-reinforced elastomer in high efficiency. Still another object thereof is to provide a pneumatic tire having excellent performances by the use of said resin-eniforced elastomer.

The present invention which has been developed for the above purposes provides a resin-reinforced elestomer which comprises a rubber and a polyoletin as principal components and which is characterized in that (1) the average particle diameter of the polyoletin is at most 1 µm, and (2) the rubber and the polyoletin are linked together. In this case, it is preferable that the resin-reinforced elestomer have a constitution wherein the rubber and the polyoletin are linked toesther via a brinding apent (c).

In addition, the present invention provides a process for producing a resin-reinforced elastomer which comprises the steps of melt kneading a polyolefin (a), a No.1 rubber (b) and a binding agent (e) to prepare a thermoplastic composition (A) having a constitution wherein the No.1 rubber (b) is dispersed in the matrix of the polyolefin (a); and adding a No.2 rubber (B) thereto, while melt kneading the resultant mixture to subject the matrix of said thermoplastic composition (A) to phase transition and disperse the polyolefin (a) in the rubber phase.

Moreover the present invention provides a pneumatic fite which comprises the resin-reinforced elastomer used in its trade, said resin-reinforced elastomer being prepared by blending eald thermoplastic composition (A) and No.2 rubber (B) so that the content of the polyolefin (a) falls within the range of 1 to 40 parts by weight based on 100 parts by weight of the total sum of the No.1 rubber (b) and the No.2 rubber (B). It is prelerable that the average particle diameter of the polyolefin in the above-mentioned resin-reinforced elastomer be at most 1 µm and that the aforesaid polyolefin be at least one member selected from the group consisting of polyothylene, polypropylene and sthylene-propylene copolymer.

The resin-reinforced elastomer according to the present invention comprises a rubber and a polyolefin (a) as

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principal components. The polyolefin to be used as the component (a) having a high melting point or a high Vicat softening point causes inferior processability, when formed into an elastoner, whereas that having a low melting point or a low Vicat softening point gives rise to inferior strength and heart ensistance, when formed into an elastoner. Printerably, the polyolefin as the component (a) has a melting point in the range of 80 to 250°C from the viewpoint of the balance among processability, strength and heat resistance, also has a Vicat softening point of at least 50°C, especially in the range of 50 to 20°0°C, and further has a meltit flow index (MPF) in the range of 0.2 to 100°, 10° min.

As such a polyclefin, use is preferably made of a homopolymer or a copolymer each of an olefin having 2 to 8 carbon atoms, a copolymer of an olefin having 2 to 8 carbon atoms and acrylic acid or an eater thereof, a copolymer of an olefin having 2 to 8 carbon atoms and an aromatic vinyl compound such as styrene, chlorosylene and a "nethigstyrene", a copolymer of an olefin having 2 to 8 carbon atoms and vinyl acetate and a copolymer of an olefin having 2 to 8 carbon atoms and vinyl acetate and a copolymer of an olefin having 2 to 8 carbon atoms and a vinylsiane compound.

The polyolefin as the component (a) oen be selected for use from a variety of polyolefine without specific limitation, and is exemplified by polyethylene base, polypropylene base, a higher polyolefin, a polyolefin comprising a monomer having at least 4 carbon atoms; a such as polybratene base, polypentene base include high density polyethylene (HDPE), in density polyethylene (LDPE) and limes rive density polyethylene (LDPE) and limes rive density polyethylene (LDPE), a lange-tead polyethylene such as chiorinated polyethylene, to the limes rive density polyethylene and a copolymer of estipliene and monomer other than olefin such as strytene-vinyl scatter copolymer, ethylene-active and estipliene and such as chiorinated polyethylene and estipliene and scatter copolymer, ethylene-exprise add expolymer, ethylene-exprise copolymer, ethylene-exprise and estipliene-vinyl-intoloxyleilane copolymer, as the polypropylene base include propylene homopolymer, a copolymer of propylene and a monomer other than olefin such as propylene -divrene copolymer and estylene-polyprene block copolymer and estylene-propylene and estylene-polyprene block copolymer and estylene-polyprene block copolymer and estylene-polyprene polyprene polyprene and estylene-propylene and express the purpose of use of the reservice increade also exercing to the purpose of use of the reservice increade also exercing to the purpose of use of the reservice increade also exercing to the purpose of use of the reservice formed establishmen. In particular, polypethylene base, polypropylene base and ethylene-propylene copolymer are preferable from the viewpoint of competibility with rubber and industrialization with at least one other.

On the other hand, the rubber component in the resin -reinforced elestomer comprises a No.1 rubber as the component (i) and 8 No.2 rubber as the component (ii) is rubber (ii) and 8 No.2 rubber as the component (iii) is rubber (iii) and in the form of rubber at room temporature, and has a glass transition temperature of 0°C or lower, particularly preferably x20°C or lower. When the glass transition temperature thereof is higher than 0°C, there is a fear that the function as the rubber is less and to be exhibited at low temperatures. The Mooney viscosity (ML₁₊₄, 100°C) of the No.1 rubber is in the range of usually 20 to 150, preferably 30 to 80. The Mooney viscosity thereof less than 20 leads to deterioration of physical properties when made into vulcanized ubber as the case may be, whereas the Mooney viscosity thereof one than 150 sometimes results in worsened processability thereof more than 150 sometimes results in worsened processability thereof more than 150 sometimes results in worsened processability thereof more than 150 sometimes results in worsened processability.

Examples of the No.1 rubber as the component(b) include diene base rubber such as natural rubber (NR), Isoprens rubber (IR), butadiene rubber (BR), 1,2-polybutadiene (1,2-BR), styrene-butadiene rubber (SBR), isoprene rubber in liquid form, butadiene rubber in liquid form, styrene-butadiene rubber in liquid form, nitrile rubber (NBR), chloroprene rubber (CR), nitrile-isoprene rubber, nitrile-chloroprene rubber, styrene-chloroprene rubber, styrene-isoprene rubber, vinylpyridine-butadiene rubber, butyl rubber (IIR), chlorinated butyl rubber, brominated butyl rubber, carboxylated styrene-butadiene rubber, carboxylated nitrile-butadiene rubber, styrene-butadiene block copolymer, styrene-isoprene block copolymer, carboxylated styrene-butadiene block copolymer and carboxylated styrene-isoprene block copolymer, polyclefin base rubber such as chlorinated polyethylene, chlorosulfonated polyethylene (CSM), ethylene-vinyl acetate copolymer, ethylene-propylene rubber (EPM), ethylene -propylene-diene copolymer (EPDM), ethylenebutene rubber and ethylene-butene-diene copolymer, rubber having the main chain of polymethylene type such as polychlorotrifluoro-ethylene, acrylic base rubber (ACM), ethylene-acrylic base rubber, fluorinated rubber and hydrogenated nitrile-butadiene rubber; rubber having oxygen atom in the main chain such as polyester-based thermoplastic rubber, epichlorohydrin rubber (CO), ethylene oxide-epichlorohydrin rubber (ECO), ethylene oxide-epichlorohydrinallyl glycidyl ether copolymer and propylene oxide-allyl glycidyl ether copolymer; silicone rubber such as polyphenylsiloxane, polydimethylsiloxane, polymethylethylsiloxane and polymethylbutyl siloxane; ruber having nitrogen atom and oxygen atom in addition to carbon atom in the main chain such as polyamie-based thermoplastic rubber, nitroso rubber, polyester urethane and polyether urethane; and the like rubbers. Use can be made of derivatives of these rubbers such as epoxy-modified, silane -modified, or maleate-modified rubber. Any of these rubbers may be used alone or in combination with at least one other.

The No 2 rubber to be used as the component (B) may be selected from the group same as the group from which said No.1 rubber is selected for use. Any of the No.2 rubbers may be used alone or in combination with at least one other. It is preferable in the resin-reinforced elastomer according to the present invention, that the aforesaid rubber and polyolelin be bonded, especially was binding agent (c). The binding agent as the component (c) may be selected for use from those that are usually employed as a binding agent for high polymers without specific limitation, and is ex-

emplified by silens coupling agents, titanette coupling agents, novolat-based alkythenol-formatdehyde initial condensate, resol-based alkythenol-formatdehyde initial condensate, novolat-based phenol-formatdehyde initial condensate, resol-based phenol-formatdehyde initial condensate, unsaturated carboxytic sold, derivatives thereof and organic periodics. Of these, silens coupling agents are preferable from the advantage that they are less apt to cause the component (a) or (b) to get, and besides capable of timely bonding to the interfaces of these components.

Specific examples of the silene coupling agent include viryltrimethoxysilane, virytriscoysilane, virytris (β-methoxysilane, virytriscathoxysilane, virytriscathoxysilane, y-glycidoxypropyrinethydimethoxysilane, y-glycidoxypropyrinethydimethoxysilane, y-glycidoxypropylethydimethoxysilane, N-β-aminotypylanipoxypropylethydiathoxysilane, N-β-aminotypylanipoxypropylethydiathoxysilane, N-β-aminotypylanipoxypropylethydiathoxysilane, N-β-aminotypylanipoxypylane, N-β-aminotypylanipoxypylane, N-β-aminotypylanipoxypilane, N-β-aminotypylane, N-β

Examples of the binding agent other than the silane coupling agent include titanate-based coupling agent such as isopropylisostearoyl (itanate, isopropylisostearoyl titanate, isopropylisostearoyl titanate, bis (dioctylpyrophosphate) oxyacetate titanate, isopropylitrioctanoyl titanate,

In the case where the silane coupling agent is used as the component (c), it may be used in combination with an organic peroxide. It is thought that by the combinational use of the silane coupling agent and the organic peroxide, a radical is formed on the molecular chain of the polydeline as the component (a) and is reacted with the silane coupling agent, whereby the reaction between the component (a) and the silane coupling agent is accelerated. The amount of the organic peroxide to be used is in the range of preferably 0.01 to 1.0 parts by weight based on 100 parts by weight of the component (a). When natural rubber or polysoprener (ubbe relaving lacoprene structure) is used as the component (b), the organic peroxide need not be used, since the aforesaid reaction takes place by shearing at the time of kneeding.

The organic perceive is preferably used which has a half-life temperature for one minite in the range of a meltkneading temperature to a temperature higher than the same by 30°C, approx. That is to say, the organic per

There are available a variety of organic peroxides, which are specifically exemplified preferably by peroxyketal active duct as 1,1-di(t-buty)peroxy)-3,3,5-timethylcyclohexane, 1,1-di(t-buty)-peroxy) cyclohexane, 2,2-di(t-buty)peroxy) ybutane; 4-di(t-buty)peroxy) valerianic acid n-butyl seter; and 2,2-buts(4-di-buty)peroxyc)clohexane)propane; and alkyl perester such as 2,2-timethylperityl peroxyneodecancete; 4-butyl peroxyleocate; butyl peroxyneodecancete; 1-butyl peroxyneo

The foregoing resin-reinforced elastomer has the structure in which the rubber component comprising the No.1 rubber(b) and the No.2 rubber (B) forms matrix, and the component(a) is dispersed therein in the form of island, and the aforeadic component(a) and (b) are bonded to each other on the interface, especially via a binding agent(c), thereby enabling the reinforcing filter to fully manifest the working effect thereof. The component(a) in the resin-forced, elastomer is uniformly dispersed as fine particles having an average particle dameter of at most 1 µm, preferably 0.01 to 1 µm.

With regard to the content of each of the components in the resin-reinforced elastomer, the content of the component (b) is preferably 20 to 250, more preferably 50 to 200 parts by weight per 100 parts by weight of the component (a). The content of the component (b), when more than 250 parts by weight of 100 parts by weight of the component (a) makes it difficult for the component (a) to form matrix in the thermoplastic composition as described hereinafter, thus causing difficulty in pelletizing and worse workability, whereas the content thereof, when less than 20 parts by weight per 100 parts by weight of the component (a) brigs about the tendency that stificient phase transition is made impossible to carry out at the time of melt kneading with the component (B), thereby worsoning the dispersion of the component.

The content of the component (c), though not being particularly limited, is preferably 0.1 to 2.0, more preferably 0.2 to 1 part by weight per 100 parts by weight of the component (a). The content of the component (c), when less than 0.1 part by weight per 100 parts by weight of the component (a), makes it difficult to obtain a thermoplastic composition having high attenght (the structure in which the componene(b) is dispersed in the matrix of the component(a) as described heroinalter j, whereas the content thereof, when more than 2.0 parts by weight per 100 parts by weight p

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the components (a) and (b). As a result, a thermoplastic composition is obtained which has an MFI in the range of 0.01 to 50 g /10min. (190°C, 5 0 kg) and a density of 0.900 to 1.200 g /cm². An MFI of more than 50 g /10min is undevorable because of decreased mechanical strength; whereas an MFI of less than 0.01g /10min is also unfavorable because of lowered fluidity and difficulty in handling. In addition, a density exceeding 1.200 g/cm³ is unfavorable, since the effect lowards lightweightness is decreased by a decrease in difference in density between ordinary rubber-reinforcing agents such as carbon black, inorganic filters, etc. and the thermoplastic composition.

Preferably, the content of the component (a) is in the range of 1 to 40 pents by weight per 100 parts by weight of the vulcanizable rubber component (content of the No.1 rubber as the component (b) and the No.2 rubber as the component (content of the component (a) of less than 1 part by weight lowers the modulus, strength, wear resistance, fatigue resistance and the like of the vulcanizate, and causes the tendency to decrease the effect towards lightweightness, whereas the content thereof of more than 40 parts by weight causes the tendency to decrease the rubber elasticity. The content of the component (a) is preferably 4 to 60 parts by weight.

The process for producing the resin-reinforced elastomer of the present invention is not specifically initiad, provided that said process is expelle of producing the rein-reinforced elastomer having the foregoing properties. According to the process of the present invention as described hereunder, the desirable resin-reinforced elastomer can efficiently be produced.

In the process of the present invention, the thermoplastic composition (A) is firstly prepared by melt kneading 100 parts by weight of the polyoletina site component (a), the No.1 rubber as the component(b) in an amount of preferably 20 to 250, more preferably 50 to 200 parts by weight, and the binding agent as the component(c) in an amount of preferably 0.1 to 2.0, more preferably 0.2 to 1.0 part by weight. The resultant thermoplastic composition (A), which has an MFI of 0.01 to 50g/10min. (190°C, 5.0 g.), approx. and a density of 0.900 to 1.200 g / om², is of the structure in which the component(b) is dispersed in the matrix of the component(a).

The form or shape of the thermoplastic composition (A) is not specifically limited, but is preferably in the form or pellet. The reason is that a pellet protein tacilitate the transfer and handling of itself as well as uniform kneading with the component(B) as described hereinafter, enables carbon black and the like to be uniformly and readily blended, kneaded and dispersed, and makes it possible to easily obtain the resin-reinforced elastomer wherein fine particles of the oldfin far unformly dispersed therein.

Subsequently, the objective resin-reinfored elastomer is produced by adding the No.2 rubber as the component (S) to the thermoplastic composition (A) thus obtained so that the component(a) amounts preferably to 1 to 40 parts by weight based on 100 parts by weight of the total of the components (b) and (8), and melt kneeding the resultant mixture to subject the matrix of the thermoplastic composition (A) to phase transition and to ultramy disperse the components (b).

In this case, by melt keeding the mixture at a temperature higher than the melting point of the component(a) in the intermoplastic composition (A), preferably by at least 5°C, more preferably by at least 10°C, the component(a) which has formed the matrix in the thermoplastic composition (A) is subjected to phase transition with the result that the component(a) is subjected to intercretispersion in the marktic consisting of the No.1 rubber and the No.2 rubber. A temperature lower than the melting point of the component(a) leads to difficulty in the cocurrence of the phase transition and in sufficient diseasesion.

The resin-reinforced elastomer is prepared, for example, by the following steps:

- Step (1): melt kneeding the components (a) and (c) for 3 to 5 minutes at a temperature higher than the melting point of the component (a), preferably by at least 10°C to cause reaction.
- Step (2): melt kneading the reaction product in the step (1) and the components (b) for 3 to 5 minutes at a temperature higher than the melting point of the component (a), preferably by at least 10°C.
- Step (3): extruding the melt kneaded product in the step (2) at a temperature higher than the melting point of the component (a), and pelletzing the extruded product at a temperature lower than the melting point of the component (a).
 - Step (4): melt kneading the pelletized thermoplastic comosition (A) obtained in the step (3) and the components (B) at a temperature higher than the melting point of the component (a) preferably by at least 10°C.

In addition, the Stops (1) and (2) may be carried out simultaneously or in reverse order. That is to say, the components (a) and (b) may be most blended in Step (1), and then the institute may be nearbed with the component (c) in Step (2). Alternatively, the components (a), (b) and (c) may be mad blended simultaneously to proceed with the reaction.

The foregoing melt blending can be carried out by means of machinery or equipment which is usually used for knowlaing a resin or rubber. Examples thoreof include Benbury mixers, kneaders, kneading extruders, open rolls, singlescrew kneaders and twin-screw kneaders.

The resin-reinforced elastomer may further be blended with an ordinary short fiber, which is exemplified by nylon short fiber, polyester short fiber, aramid short fiber, cotton short fiber, vinylon short fiber, rayon short fiber natural

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cellulose short fiber and acrylic short fiber

In the kneading step for the component(B), a chemical agent may be added and kneaded as necessary, which is usually employed in the rubber industry and is exemptified by vulcanizing agents, vulcanizing aids, filters, age resistors, process oils, zino oxide and stearic acid.

The amount of a vulcanizing agent to be added therein is not specifically limited, but is in the range of preferably 0.1 to 5.0, particularly preferably 0.5 to 3.0 parts by weight based on 100 parts by weight of the rubber components. The amount of a vulcanizing aid, when being added, is in the range of preferably 0.01 to 2.0, particularly preferably 0.1 to 1.0 parts by weight based on 100 parts by weight of the rubber components.

Examples of the vulcanizing agents include well-known vulcanizing agents such as sulfur, organic peroxides, resing vulcanizing agents and metal oxides such as magnesium oxide.

The vulcanizing aid is selected from well-known vulcanizing aids such as aldehydes, ammonia, amines, guanidine, thicurea, thiazole, thiurams, dithiccarbamate and xanthate.

Examples of the fillers include reinforcing agent such as carbon black of various types, white carbon, activated calcium carbonate, ultrafine particulate magnesium silicate, high impact polystyrene, phenolic resins, lignins, modified melamine resin, cumarone indene resins and petroleum resins, calcium carbonate, basic magnesium carbonate, clay, litherge, diatomaceous earth, regenerated rubber and powdery rubber.

Examples of the age resistor include amine-ketones, imidazoles, amines, phenois, sulfur base and phosphorus

The usable process oil may be any of aromatic base, naphthenic base and paraffinic base.

The vulcanization temperature of the resin-reinforced elastomer may be the vulcanization temperature of ordinary rubber and is preferably in the range of 100 to 190°C.

The foregoing resin-reinforced elastomer is widely and effectively used for the external members of tires such as tire tread and tire side-wall, internal members of tires such as carcass, bead and chafer, and rubber members for Industrial products auch as hosse, bettis, rubber rolls and rubber crawlers.

In the following, some description will be given of the pneumatic tire using the resin-reinforced elastomer of the present invention.

In the treed of the pneumatic thre according to the present invention, use is made of the foregoing resin-reinforced element within is produced by blending the aforesaid thempolastic composition(A) and the No.2 rubber as the component(B) and make three the resultant mixture, said thempolastic composition(A) having the structure in which the component(b) is dispersed in the matrix of the component(b) which matrix is formed by melt kneading the No.1 rubber as the component(b) in an amount of preferably 20 to 250, more preferably 50 to 200 parts by weight, and the binding agent as the component(b) in an amount of preferably 0.1 to 2.0, more preferably 0.2 to 1.0 part by weight each based on 100 parts by weight of the polyolefin (a). The form of the thermoplastic composition (A) is particularly preferably pell from the aspect of workability.

The foregoing realn-reinforced elastomer is capable of manifesting the effect as the reinforcing filler inrough the constitution in which the rubber component consisting of the No.1 rubber as the component(b) and the No.2 rubber as the component(b) forms the matrix; the polyelefin (a) in the form of island is dispersed in the rubber component in an amount of 1 to 40, preferably 2 to 30 parts by weight based on 100 parts by weight to the rubber component; and the components (a) and (b) are bonded to each other on the interface thereof via the binding agent. The polyelefin fine as the component (a) in the restin-reinforced elastomer is uniformly dispersed as fine particles having an average particle diameter of preferably all one titum, more preferably 0.01 to turn.

The polyolefin as the component(a) may be properly selected for use from those described hereinbefore, and is particularly preferably polyethylene, polypropylene or ethylene-propylene copolymer. Any of these polyolefins may be used alone or in combination with at least one other.

The types of the Nos. 1 & 2 rubbers are as variously exemplified before and are preferably natural rubber (NF), isoprene rubber (IR), butselfene nubber (IRF), 12-polybutselinen (12-BR) or styrene butselfien en blood (IRF). Profer (NR), natural rubber or isoprene rubber occupies at loast 40% by weight of the total amount of the rubber components (No rubber (b) bits No 2 rubber (IRF).

Moreover, the resin-reinforced elastomer may be blended with carbon black as a filler in an amount of usually 30 to 70 parts by weight based on 100 parts by weight of the total amount of the rubber components.

In the following, the present invention will be described in more detail with reference to comparative examples and working examples, which however shall not limit the present invention thereto.

Various measurements were performed as follows:

(1) Evaluation method for the thermoplastic composition(A)

Possibility of pelletization

Pelletizable composition was marked with O, and non-pelletizable composition was marked with X.

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@Melt Flow Index (MFI)

The amount of mo'ten resin passed through an orifice having an inside diameter of 2 mm and a length of 8 mm at a load of 5000g at 190°C was expressed in g /10min. as MFI.

③Density

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Measurements were made of density according to JIS K7112 by means of a density gradient tube.

(2) Evaluation method for the resin-reinforced elastomer

Tensile stress, tensile strength and elongation at break

According to JIS K6251, measurements were made of the tensile stress, tensile strength and elongation at the strength and advantage at 100% and 300% in both parallel and perpendicular directions against sheet discharge direction by punching sheet into test specimen according to JIS dumb-bell Shep Pattern No.3.

@Hardness

Measurements were made of hardness according to JIS K6251 by type A.

Abrasion resistance

Evaluations were made of abrasion resistance according to JIS K6264 by measuring Lambourn abrasion using indexes based on an Index in Comparative Example 1 as 100.

②Fatigue resistance

Evaluations were made of fatigue resistance using a constant-elongation-type fatigue feating machine (produced by Yesuda Seliki Selasukan b) by applying a load of 500g and a constant strain of 100% repeatedly to determine the number of times until the test specimen was cut off. (@Particle diameter.

Measurements were made of the particle diameters of the polyolefin dispersed in the rubber matrix to determine the average particle diameter by the use of electrophotographs of the vulcanized resin-reinforced elastomer.

(3) Evaluation method for the performance of test tire.

(1) Cut-through resistance and chipping resistance.

Each of tires was mounted on a 10 ton truck and was subjected to running test mainty on rough and unpawed roads at an internal pressure of 0.71 MPa until the wom out condition (maximum depth of residual grove being at most 3 mm.).

Cut-through resistance was determined by peeling the tire tread and measuring the number of cuts which reached the belt through the tread.

Chipping resistance was determined by measuring the number of chips being at least 1 cm² in size within the surface of tire tread in contact with ground.

Durability against heat generation

According to the durability test procedure prescribed in JIS D4230-1968 Table 5.5, the durability against heat generation was evaluated by measuring the running range which causes tire breakage to the extent of being incapable of running, using indexes based on an index in Comparative Example T-2 as 100. The durability against heat generation excels with an increase in said index.

(3) Abrasion resistance

Each of tires was mounted on a truck and was subjected to continuous running test for a distance of 0,000 km, when an evaluation was made of the abrasion resistance thereof by measuring the tread groove depth, using indexes based on an index in Comparative Example T-2 as 100. The abrasion resistance excels with an increase in said index.

@Processability

Measurements were made of the viscosity of unvulcanized rubber kneaded for use in the aforesaid tire tread under the conditions of 100°C and a shear rate of 10 see? by the use of a processability testing machine (produced by Monsanto Co., Ltd. under the trade name* MPT*) to evaluate processability, using indexes based on an index in Comparative Example T-2 as 100. The processability excels with an increase in said index.

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[Preparation of thermoplastic composition (A)]

Preparation Example 1

Preparation of Sample 1:

There were used a high density polyethylene (HDPE produced by Maruzen Polymer Co.,Ltd. under the trade name "Chemilates HD 3070"; density of 0.952 g/cm², MFI of 30 g/10 min, melting point of 130") as the component (a). NR (SNR-L) as the component(b) and y-methacryloxypropythimethoxysilane as the component(c). At first, 100 parts by weight of the component(c) were melt kneaded at 140"C for 31 minutes with a B-type Barbury mixer with 1.72 capacity, and then the melt kneaded product and 150 parts by yeight of the component (b), each based on 100 parts by weight of the component (d), were melt kneaded at 150"C for 3 minutes with a B-type Barbury mixer with 1.72 capacity in the same manner as before. The kneaded product and subject to the same manner as before. The kneaded product subject of the component (b), and a subject of the component (b) and the same manner as before. The kneaded product subject of the same manner as before. The kneaded product was dumped at 170"C and was pelletized at a die temperature of 160"C. The objective product had a density of 0.934g /cm², MFI of 2.0 g 1/0 min, and chemicals composition and physical proporties as given in Table 1.

Preparation Example 2

Preparation of Sample 2

The procedure in Preparation Example 1 was repeated to prepare Sample 2 and pelletize the same except that the kineading was carried out by the use of the component (a) consisting of polypropylene(PP, produced by Ube Industries Ltd. under the trade name? Ube Polypro 19969 3 (ensity of 9.05 g/cm²), MFI of 3.0 g/10 min., melting point of 148°C) at 150°C for 3 minutes. The objective product had a density of 0.913g/cm², MFI of 1.90 g/10 min. and a chemical composition and previously processing and processing processing and previously processing processing and previously processing and processing processing and processing processing and processing pro

Preparation Example 3

Preparation of Sample 3

The procedure in Preparation Example 2 was repeated to prepare Sample 3 and polletize the same except that the use of the component(c) was omitted. The resultant product had a density of 0.912g/cm³, MFI of 2.5 g /10 min, and a chemical composition and physical properties as given in Table 1.

Preparation Example 4

Preparation of Sample 4

The procedure in Preparation Example 2 was repeated to prepare Sample 4 except that the melt kneeding was carried out by the use of the component(a) consisting of ultrahigh molecular polyethylene(HMPE, produced by Mitsul Petrochemical industries, Ltd. under the trade name* Hizex-Million 340M*, density of 0,939 g /cm², melting point of 198°C). However, pelletizing was impossible. The chemical composition and physical properties of the non-palletized product are given in Table 1.

45 Preparation Example 5

Preparation of Sample 5

The procedure in Preparation Example 2 was repeated to prepare Sample 5 except that the melt kneading was carried out by the use of 500 parts by weight of NR as the component (b): However, pelletizing was impossible. The chemical composition and physical properties of the non-paletized product are given in Table 1.

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Sample No.	1	2	3	4	5
Thermoplastic					
composition (A)					
Component (a) (wt.parts)	HDPE (100)	PP (100)	PP (100)	HMPE (100)	PP (100)
Component (b) (wt.parts)	NR (130)	NR (130)	NR (130)	NR (130)	NR (500)
Component (c) (wt.part)	MPMS (1) MPMS (1)	MPMS (1)	1 :	MPMS (1)	MPMS (1)
Palletizing					
Possible ()	0	0	0		
Impossible x				×	×
MFI (g/10min.)	2.0	1.9	2.5	< 0.01	< 0.01
Density (g/cm*)	0.934	0.913	0.912	1	1

[Remarks] MPMS : 7 -methacryloxypropyltrimethoxysilane

[Preparation of Resin-reinforced Elastomer]

Examples 1 to 4

In a 250 mt plastomill for laboratory use which had been set to 120°C and 68 r.p.m. were placed a sample of the there proposed to the proposed composition (A) and NR as the component (B) in the blending formulation as shown in Table 2; followed by mastication for 30 seconds. Then the resultant mixture was further blended with carbon black (produced by Mit-

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subish Chemical Industries Ltd, under the trade name ISAF, average particle diameter of 21 µm, DBP oil adsorption of 117mf /1090, zinc oxide, stearic acid and an age resistor (N-phenyl-N-isoprophy-henylonodiamino, trade name/810NA*), followed by kneading for 3 minutes. The the dumped product, the temperature of which was 180 to 170°C, was blended with a vulcenization accelerator DM (dibenzothiazi) suitide), a vulcenization accelerator DM (dibenzothiazi), a vulcenization accelerator DM (dibenzothiazion), a vulcenization accelerator DM (dibenzothiazion

Comparative Example 1

The procedure in Example 1 was repeated to prepare a resin-reinforced elastomer except that polypropylene resin was used in place of the thermoplastic composition (A) in accordance with the formulations as shown in Table 2. Evaluations were added to the physical properties of the resultant resin-reinforced elastomer. The results are given in Table 2. This elastomer was inferior to that in the examples in terms of albrasion resistance as well as altique resistance.

Comparative Example 2

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The procedure in Example 1 was repeated to prepare a resin-reinforced elastomer except that the amounts of the thermoplastic composition (A) (Sample 2) and NR as the component(B) were altered in accordance with the formulation as shown in Table 2. Evaluations were made of the physical properties of the resultant resin-reinforced elastomer. The results are given in Table 2. In spite of its very high modulus as compared with the examples, this elastomer had a low tensile strength and elongation and was inforior to the elastomers in examples in terms of abrasion resistance and fatigue resistance, without forming complete sea-sland constitution.

25 Comparative Examples 3 to 5

The procedure in Example 1 was repeated to prepare resin-teriforced elastomers except that the amounts of the thermoplastic composition (A) I samples 3 to 5) and NR as the component (B) were altered in accordance with the formulation as shown in Table 2. Evaluations were made of the physical properties of the resultant resin-reinforced elastomers. The results are given in Table 2. Those elastomers were inferior to those in the examples in terms of abrasion resistance as well as fatigue resistance.

vulcanization

carbon black (40), zinc oxide (3), stearic acid (2), age resistor-810NA (1), accelerator DM (0.6), vulcanization accelerator D (0.2), sulfur (1.5)

Thermoplastic Composition (A Sample No. 1 2 3 4 1 2 3 4 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5			H	Table	7					
1 2 3 4 1 2 3 4 1 2 3 4 4 1 2 3 4 4 4 5 5 4 4 5 5 4 4			Exampl	a No.		Ü	omparat	ive Bxa	on elon	
1, 10 4 2 2 2 - 2 3 4		-	7	ю	4	-		,		
1, 10. 1 2 2 2 - 2 3 4	Thermoplastic Composition (×	L	L	L			,	-	6
1341 46 23 46 69 - 15 53 23 23 24 24 25 25 25 25 25 25	Sample No.	-	7	8	2	_ '	٠	,	•	
143410	(part/s by weight	_	23	46	9		• :	° ;	-	<u>.</u>
1960 1974 87 74 61 100 35 74 74 100 10		\perp			١	1	=	23	23	9
100 14 174 174 11 100 35 74 74 100		1	,	1	,	2	,	1	ı	1
100 100	NR (part/s by weight)		87	74	- 19	e e	¥	;		
1982 3.7 3.5 4.9 6.1 3.9 19.6 3.7 3.7 3.7 3.8 4.9 6.1 3.9 19.6 3.7 3.7 3.4 4.5 5.7 3.3 15.4 3.3 3.4 4.5 5.7 3.3 15.4 3.3 3.4 4.5 5.7 3.3 15.4 3.3 3.4 4.5 5.7 3.3 15.4 3.3 3.4 4.5 5.7 3.3 15.4 3.3 3.4 4.5 5.7 3.3 15.4 3.3 3.4 4.5 5.7 3.3 15.4 3.3 3.4 4.5 5.7 3.5 5.5	olyolefin amount per 100	L					;		•	20
190 3.7 3.5 4.9 6.1 3.9 18.6 3.7 3.7 3.7 3.8	parts by weight of rubber	-,-								
Pa 3.7 3.5 4.9 6.1 3.9 18.6 3.7 3.7 3.7 3.8 4.9 6.1 3.9 18.6 3.7 3.7 3.7 3.8 4.9 6.1 3.9 18.6 3.7 3.7 3.7 3.8 3.9 18.6 3.9	omponent (part/s by weight)	_	2	20	9	5	5	;		
1 3.7 3.5 4.9 6.1 3.9 18.6 3.7 3.7 3.7 3.4 3.9 18.6 3.7 3.4	Physical Properties						2	2	٩	2
1 3.6 3.3 4.5 5.7 3.3 15.4 3.3 3.4 1 1 1 1 1 1 1 1 1	MPa)	3.7	3.5	4.9	6.1	3.9	18.6	,	,	
11.8 13.6 16.4 20.2 14.5 13.6 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8 13.7 13.8	Т	3.6	3.3	4.5	5.7	3.3	15.4		; ;	, ,
11.6 13.3 15.5 19.1 13.6	300% tensile stress (MPa)	11.8	13.6	16.4	20.2	14.5	,	; ;	;	2.3
30.1 28.3 28.4 28.2 27.8 23.1 13.3 13.7 400 550 28.4 28.6 28.1 27.5 22.5 27.5 25.5 400 550 500 450 540 170 550 480 49 73 72 78 82 73 75 55 500 49 74 75 75 76 80 100 87 103 85 75 75 75 75 75 75 75	T	11.6	13.3	7 2	- 0	: ;	ı	2	P. 4	
1	Pastle strongth (mm.)		?	2	-	2	1 -	13.3	13.7	12.9
1	(BAN) III GIR TO GETTER	20.	28.3	28.8	28.2	27.8	23.1	27.9	25.8	27.2
640 550 500 450 540 150 550 480 440 440 440 540 170 550 480 440	7	29.6	28.4	28.6	28.1	27.5	22.5	27.5	25.5	9,0
14 640 560 510 460 540 170 550 500 49 13 72 78 82 73 95 72 71 7 14 15 105 102 100 87 103 85 10 15 0.38 0.39 0.32 0.51 0.72 1.21 0 0.54 0.38 15 0.38 0.38 0.37 0.72 1.21 0 0.64 30.10 1 1 1 1 1 Parpandicular Other Components Inspector	Clongation (%)	640	550	200	450	540	150	550	480	7 0
13 12 78 82 13 95 72 70 13 10 10 10 10 10 10 10 10 10 10 10 10 10	7	640	260	510	460	540	170	550	200	2 9
Ders 65,000 95,000 23,000 15,000 00,000< 1,000 52,000< 1,000 35,000 13,000 15,000 10,000< 1,000 52,000< 1,000 35,000 13,000 13,000 10,100 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000 1,000 35,000 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000< 1,000 52,000	Hardness (JIS-A)		72	78	82	73	95	23	3 5	7.7
10 10 10 10 10 10 10 10 10 10 10 10 10 1	Abrasion resistance (index)	Ξ	113	105	102	100	. 48	103	- 4	- 5
0.38 0.32 0.51 0.72 1.21 - 0.64 30.10 1 :Porpondicular Other components [part/s by margate]	atigue resistance:numbers	85,000	000'96	23,000		20,000	< 1,000		1.000	35 000
:parallel 1 :perpendicular Other components (part/s by notable)	rticle diameter (m)	0.38	0.32	0.51	0.72	1.21	,		30.10	
	_	T : per	pendicu	lar	Other	Compone	ints (par	t/e hv	140,00	- 1

[Preparation of test tires and evaluation of performance]

In accordance with the blending formulation as shown in Table 3, tires of the size 10.00R2014PR were prepared by a conventional method by using, in the tire tread, a resin -reinforced elastomer which had been prepared from the compounding ingredients including the thermoplastic elastomer (A), component (B), carbon black (HAF), aroma oil and the like. Thus the performances of the resultant tires were examined.

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Comparative Examples T-1 to -T-4

Test tree were prepared by using, in the tire treads, rubber compositions wherein the blending amounts of carbon black (HAF) which is the typical means for mcdifying the characteristics of tire tread were varied from 40 to 55 parts by weight. Thus the performances of the resultant tires were examined. The results are given in Table 3.

Comparative Example T-5

Test tires were prepared in the same manner as in Comparative Example T-4 except that the thermoplastic composition (Sample 1) was blended so that the amount of the polydefin was made to 0.5 part by weight based on 10.0 parts by weight of the whole rubber components. The results are given in Table 3. As can be seen from the table to improvement effect is recognized against Comparative Example T-4 in which the amount of carbon black was same as herein, but no polydefin was used.

5 Examples T-1 to -T-5

Test tires were prepared in the same manner as in Comparative Example T-4 except that the thermoplastic composition (Semple 1) was blanded so that the amount of the polyodefin was made to 1.5 to 10 parts by weight based on 100 parts by weight of the whole nubber components. The results are given in Table 3. As can be seen from the table, these lest tires are remarkably excellent in both durability against heat generation and abrasion resistance and also in axtituation processability as compared with Comparative Example T-1 to 1-4.

Examples T-6 to -T-7

Test tires were prepared in the same manner as in Comparative Example T-4 except that the thermoplastic composition (Sample 1) was blended so that the amount of the polyolefin was made to 15 to 18 parts by weight the whole rubber components. The results are given in Table 3. As can be seen from the table, these test tires are excellent in all respects including durability against heat generation, abraidon resistance that and cut -through resistance as compared with Comparative Example T-1 in which the extrusion processability is almost the same as those examples.

Examples T-8

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Test tires were prepared in the same manner as in Example T-3 except that an increased amount of carbon black was added to the rubber components. Various performances of the resultant tires are given in Table 3. As can be seen from the table, these test tires are excellent in all respects including out-through resistance, durability against heat generation and abrasion resistance as compared with Comparative Example T-1.

Examples T-9

Test tires were prepared in the same manner as in Example T-3 except that use was made of the tharmoplastic composition (Sample 2) incorporated with polypropylene as the polypropylene. Marous performance of the residuant tires are given in Table 3. As can be seen from the table, these test tires are excellent in extrusion processability, durability against heat generation and abrasion resistance in a well belanced manner as compared with Comparative Examples T-1 to T-4, and are also excellent in extrusion processability and abrasion resistance as compared with Comparative Examples T-7 as described hereinsfer in which only polypropylene was added to the rubber components at the time of blending without the use of the thermoplastic composition as the component (A).

Comparative Examples T-6 to T-7

Test ties were prepared in the same manner as in Example T-3 except that only polyolefin was added to the rubber composition without the use of the thermoplasic composition as the component (A). The results are given in fable 3, As can be seen from the table, the test times in Comparative Example T-6 are inferior to those in Example T-3 wherein the thermoplastic composition (Sample 1) was used as the component (A) in regard to extrusion processability, durrability against heat generation and abrasion resistance, especially in regard to extrusion processability.

Example T-10 & Comparative Example T-8

Test tires were prepared in the same manner as in Example T-3 and Comparative Example T-4, respectively except that use was made, as the component(B), of the blend of NR and BR (produced by Japan Synthetic Ruber Co.,Lid. under the trade name *BR01*). The results are given in Table 3. As can be seen from the table, the test tires in Example T-10 are remarkably improved as compared with Comparative Example T-8 in terms of chipping resistance, cut-through resistance and abrasion resistance in spile of durability against hard generation being almost the same therebetween.

Example T-11 & Comparative Examples T-9 to T-10.

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Test tires were prepared in the same manner as in Example T-10 and Comparative Example T-6, respectively in Example T-11 & Comparative Examples T-9, except that increased amounts of carbon black and aroma oil were bended in studies tires. In Comparative Example T-10, test tires were prepared in the same manner as in Comparative Example T-6 except that use was made of increased amounts of carbon black and aroma oil along with the blend of NR and BR as the componer (IS). The results are down in table 3.

From the results of Examples T-10 à. T-11 and Comparative Examples T-9 à. T-10, it is understood that the line prepared by the use of the thermoclastic composition of the present invention are superior to those prepared without the use of said thermoclastic composition and to those prepared by exiting a polyolefin at the time of blending with regard to cut-through resistane, abrasion resistance and the like in the cese where an alteration is made of the amount of oil. It can also been seen their urboar to be used as the component (6), or an alteration is made of the amount of oil. It can also been seen their other than the time in Example T-10 are superior to those in Comparative Example T-9 with respect to chipping resistance, outflowing the resistance and abrasion resistance, outflowing the set and and brasion resistance, outflow for durability against theat generation being altmost the same therebetween; and also that the time in Example T-10 a split of the performance of time being almost the same therebetween.

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				Table	1.0	n						
		Ŭ	прага	Comparative Example	xample		L	EX	Example	10		
		P-1	T-2	7-3	T-4	T-5	7-1	4-7	F			
Thermoplastic o	Thermoplastic composition (A)	L								1	-	
	Sample 1 (ppwt)	ı	1	1	ı	1.15	3.45		9.20 11 5	9		
	Sample 2 (ppwt)	i	ı	ı	J	1	1		:		4 .	34.5
E resin	(ppwt)	ı	,	J	ŀ				1	,	,	ı
PP resin	1 1	þ	ļ		1 1	ı,	1	11	1	11]] [1
Component (B)	NR (ppwt)	100	100	100	100	1 00			1	1	,	1
	BR (ppwt)	1	J	. 1			7	ъ.	93.5	9.6	87.0	9.92
Amount of polyolefin per 100	lefin per 100					-	,	1	,	1	1	1
	ppwt of rubber component (ppwt)	•	0		c	9		,	,			
G Carbon black	(ppwt)	55	5.0	45	,	;		-	•		٥	15
Aroma oil	(hund)		;	;	3	2	40	60	40	40	40	40
Polvolefin navt	Polyolofin next(=) - 31 /	\cdot	-	-	-	-	-	-	-	-	-	-
Too at the Car	Tore dia. (# #)	,	,	,	ı	0.3	0.3	0.3	0.3	0.35	4.0	0.42
r dr rormances	S O C C C								Ī	I	Ī	
Extrusion processability	essability											
	(1ndex)	94	100	108	122	121	120	118	31.6			ļ
Chipping resistance	tance (Nos.)	-	۰	-	,	,	-			7	5	16
Cut-through resistance (Nos.)	sistance (Nos.)	ø	8	91	, c	, 4	. ;	- ;	,		•	0
Durability against heat	inst heat			!	3	3		2	n	m	~	7
den	generation (index)	94	100	108	115	118	117	115	114	100	901	8
Abrasion resistance	tance (index)	103	100	94	90	06	-	701		· :	8 ;	6
[Remarks] ppwt:pa	PPwt:part/s by weightOther components (nmmt)	othe	T COME	onente	(puert)			5		7	*	=
						. 21115	×		Local	7	ć	

components (ppwt): zinc oxide (4), stearic acid (2), paraffin wax (2), age resistor-Santoflex 13(1), vulcanization accelerator CZ(1), sulfur (2).

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			ě	Table	ტ ტ		(Continued)	Ç 0 Q			
		_	Example	_	Сомрак	Comparative	Example Comp.	Comp.	ExampleComparative	Compar	ative
						Example		Example		8	Example
		T-7	F-8	T-9	T-6	1-1	T-10	1-8	1-11	9	1.10
Thermoplastic composition (A)	on (A)					ŀ					2
Sample 1 (ppwt)	(ppwt)	41.4	1.5	1	1	1	11.5	ı		i	
. Sample 2 (ppwt)	(ppwt)	ı	i	11.5	ı	ı	1	ı	1	1 1	1 1
HDPE resin	(bbat)	ı	1	ı	s	.1	,	1	,		
	(ppwt)	ı	ı	J	1	5	1	-			0
Component (B) NR	(bbwt)	9.92	93.5	93.5	100	100	63.5	7.0	63.5	70	
88	(ppwt)	1	1	1	ı	ı	30	30	30	30	30
Amount of polyolefin per 100	r 100										
ppwt of rubber component (ppwt)	t (ppwt)	2	2	'n	S	L.	u)	•	u	-	u
Carbon black	(ppwt)	40	S	ê	40	40	40	9		2	-
Aroma oil	(ppwt)	_	-	-	-	-	-	-	L.	3 4	3
Polyolafin particle dia. (m m)	(##).	0.45	0.25	0.5	0.35	2	-	1		ŗ	
Performances									3	,	9
Extrusion processability											
2	(index)	96	9.2	114	108	95	101	106	8	128	901
Chipping resistance (N	(Nos.)	_	۰	-	-	-	7	~		, ,	
Cut-through resistance (Nos.)	fos.)	60	е,	æ	8	φ	10	28	24	۲,	,
Durability against heat					<u></u>					3	r N
generation (index)	(ndex)	95	100	116	110	116	116	119	100	102	101
Abrasion resistance (i	(index)	113	116	100	104	95	107	26	86		: 6
Remarked party	1	1			1		-			2	

|Remarks| ppwt:part/s by weight...Other components(ppwt): zinc oxide(4), stearic acid(2), paraffin wax (2), age resistor-Santoflex13(1), vulcanization accelerator $CZ\left(1\right)$, sulfur $\left(2\right)$.

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Claims

- A resin-reinforced elastomer which comprises a rubber and a polyclefin as principal components and which is characterized in that (1) the average particle clameter of the polyclefin is at most 1 µm, and (2) the rubber and the obviolefin are linked together.
- The resin-reinforced elastomer according to Claim 1, wherein the rubber and the polyolefin are linked together via a binding agent (c).
- 3. The resin-reinforced elastomer according to Claim 2, wherein the rubber in the resin-reinforced elastomer comprises a No.1 rubber (b) and a No.2 rubber (B); the content of said No.1 rubber (b) is in the range of 20 to 250 parts by weight of the polygolderin (a); the content of said binding agent (c) is in the range of 0.1 to 2.0 parts by weight based on 100 parts by weight of the polygoldin (a); and the content of said polygoldin (a) in the resin-reinforced elastomer is in the range of 1 to 40 parts by weight based on 100 parts by weight of the total sum of said No.1 rubber (b) and said No.2 rubber (B).
 - 4. The resin-reinforced elastomer according to any of the preceding claims, wherein the polyolefin particles in the resin-reinforced elastomer have an average particle diameter in the range of 0.01 to 1 µm.
- The resin-reinforced elastomer according to any of the preceding claims, wherein the polyolefin (a) is at least one
 member selected from the group consisting of polyethylene, polypropylene and ethylene-propylene copolymers.
 - 6. A process for producing a resin-reinforced eleatomer which comprises the steps of melt kneading a polyolefin (a), a No.1 rubber(b) and a binding agent(c) to prepar a thermoplastic composition (A) having a constitution wherein the No.1 rubber (b) is dispersed in the matrix of the polyolefin (a); and adding a No.2 rubber (6) thereto, which is residing the resultant mixture 1s, subject the matrix of said thermoplastic composition (A) to phase transition and disperse the polyolefin (a) in the subber phase.
- 7. The process for producing a resin-reinforced elastomer according to Claim 6, wherein the content of said No.1 or tubber (b) is in the range of 20 to 250 parts by weight based on 100 parts by weight of the polycletin (a); the content of said binding agent (c) is in the range of 0.1 to 2.0 parts by weight based on 100 parts by weight of the polycletin (a); and the content of said polycletin (a) in the resin-reinforced elastomer is in the range of 1 to 40 parts by weight based on 100 parts by weight of the total sum of said No.1 rubber (b) and said No.2 rubber (c).
- The process for producing a resin-reinforced elastomer according to Claim 6 or 7, wherein the thermoplastic composition (A) has a melt flow index (MFI) in the range of 0.010 to 50 g /10 min. and a density in the range of 0.900 to 1.200 g /cm⁻³.
- The process for producing a resin-reinforced elastomer according to any of Claims 6, 7 and 8, wherein the polyoletin
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- 10. A pneumatic tire which comprises a resin-reinforced elastomer used in its tread, said resin-reinforced clastomer being prepared by blending a No.2 ubber (8) and a thempositest composition (A) which is obtained by melt sheading a polyclefin (a), a No.1 rubber (b) and a binding agent(e) and which has a constitution wherein the No.1 rubber (b) is dispersed in the matrix of the polyclefin (a), a lost that the content of the polyclefin (a) falls within the range of 1 to 40 parts by weight based on 100 parts by weight of the total sum of the No.1 rubber (b) and the No. 2 rubber (B).
- 50 11. The pneumatic tire according to Claim 10, wherein the thermoplastic composition (A) has a chemical composition in which the content of the No.1 rubber (b) is in the range of 20 to 250 parts by weight, and the content of the binding agent (c) is in the range of 0.1 to 2.0 parts by weight each based on 100 parts by weight of the polyelinf (a).
- 12. The pneumatic tire according to Claim 10 or 11, wherein the average particle claimeter of the polyolefin (a) in the resin-reinforced elastomer is at most 1 µm.
 - 13. The pneumatic tire according to any of Claims 10, 11 and 12, wherein the polyolefin (a) is at least one member selected from the group consisting of polyethylene, polypropylene and ethylene-propylene copolymers.



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	DOCUMENTS CONSIDE	RED TO BE RELEVANT		
ategory	Citation of document with inc of relevant passa	fication, where appropriate. ges	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CL6)
	EP 0 039 240 A (JAPA LTD) 4 November 1981 * page 23, line 15 - table 3 *		1	C08L21/00 C08J3/00 //(C08L21/00, 23:02)
١	US 4 628 073 A (FISH 1986 * claims 20,37 *	ER W KEITH) 9 December	1	
	FR 2 127 217 A (KLEE October 1972 * claims *	ER COLOMBES) 13	1	
				TECHNICAL PIELDS SEARCHED (PA.CLA) COSL COSL
	The present soarch report has b	on drawn up for all claims Date or consession of the seath		
	THE HAGUE			Examiner
		9 July 1998		Humbeeck, F
X : part Y : part doc: A : tect O : non	ATEGORY OF CITED DOCUMENTS icularly relevant if taken alone cularly relevant if combined with anoth ment of the same category nological background -written disclosure mediate document	T : theory or principle E : earlier patient doc after the filling disk D : document cities L : document cred is & : member of the sa document	ument, bul publi a the apolication ir other reasons	sthed on, or

